

ACCELERATED COMMUNICATION

Pharmacological Characterization of Heterologously Expressed ATP-Gated Cation Channels (P_{2x} Purinoceptors)

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Received March 28, 1995; Accepted May 5, 1995

SUMMARY

cDNAs encoding P_{2x} purinoceptors from human bladder smooth muscle and from rat PC-12 cells were expressed in oocytes and human embryonic kidney 293 cells. Agonist potencies of 2-chloro-ATP = ATP $\geq 2'$ - and triphosphate $\geq P_1^3P_6$ -dihadenosine-5'-pentaphosphate \gg ADP prevailed for both P_{2x} purinoceptors. There were two main differences in agonist sensitivity between the two receptors. First, ATP was 10 times more potent at the receptor from the bladder (EC_{50} , 0.8 μ M) than at the receptor from PC-12 cells (EC_{50} , 8.2 μ M). Second, α - β -methylene-2',2'-disulfonic acid was a potent agonist in cells expressing the bladder smooth muscle receptor (EC_{50} , 1–3 μ M) but was ineffective in cloned P_{2x} receptor channels.

The present study addresses the third problem mentioned above, namely the possible existence in a given cell of multiple molecular species of P_{2x} receptors. Distinct cDNAs encoding ionotropic P_{2x} purinoceptors have been isolated from rat vas deferens smooth muscle and from nerve growth factor-differentiated PC-12 cells (11, 12). More recently, a human homologue of the rat vas deferens P_{2x} receptor has been obviated by making whole-cell recordings from dissociated cells, using rapid, "concentration-clamp" delivery of agonists and using the patch pipette to dialyze out cytoplasmic constituents necessary to drive G protein-coupled processes. Recently, these types of electrophysiological studies have been carried out on acutely dissociated smooth muscle cells (5, 6),

as well as cultured autonomic and central neurons and glia (7–10). These studies revealed three distinct P_{2x} purinoceptor phenotypes, i.e., an α - β -MeATP-sensitive, desensitizing inward current characteristic of smooth muscle P_{2x} receptors, an α - β -MeATP-insensitive, nondesensitizing, inward current characteristic of responses in the PC-12 pheochromocytoma cell line and superior cervical ganglion neurons, and an α - β -MeATP-sensitive, nondesensitizing, inward current that is observed in other neurons such as celiac and nodose ganglia.

The present study addresses the third problem mentioned above, namely the possible existence in a given cell of multiple molecular species of P_{2x} receptors. Distinct cDNAs encoding ionotropic P_{2x} purinoceptors have been isolated from rat vas deferens smooth muscle and from nerve growth factor-differentiated PC-12 cells (11, 12). More recently, a human homologue of the rat vas deferens P_{2x} receptor has been obviated by making whole-cell recordings from dissociated cells, using rapid, "concentration-clamp" delivery of agonists and using the patch pipette to dialyze out cytoplasmic constituents necessary to drive G protein-coupled processes. Recently, these types of electrophysiological studies have been carried out on acutely dissociated smooth muscle cells (5, 6),

cloned from human bladder smooth muscle (13). When it is expressed in oocytes or mammalian cells, activation of this purinoceptor results in robust currents due to the opening of cation-selective ion channels. This provides the opportunity to determine the pharmacological profile of individual molecular species of P_{2x} receptors by heterologous expression. The present study was undertaken to define agonist and antagonist properties of homo-oligomeric forms of the smooth muscle and properties of homo-oligomeric forms of the smooth muscle and PC-12 P_{2x} purinoceptors.

Materials and Methods

Reproduction system. Human urinary bladder P_{2x} cDNA (13), subcloned into the pBKCMV expression vector and PC-12 cDNA (13) (supplied by A. Brake and D. Julius, University of California, San Francisco), subcloned into the pBKCMV vector or the pcDNAI vector, were used for transfection of HEK 293 cells and for in vitro RNA transcription and injection into oocytes.

Lipofectin was placed in a 35-mm Petri dish containing four coverslips on which HEK 293 cells were plated (5×10^4 cells/cover slip); this medium was removed after 5–6 hr of incubation at 37° and replaced with normal culture medium, and recordings were made 12–48 hr later. Greater than 90% of cells from which recordings were made responded to ATP, but no currents in response to applied ATP (30 or 100 μ M) were observed in nontransfected ($n > 60$) or mock-transfected ($n = 22$) HEK cells. Difolliculated Xenopus oocytes were injected with 50 ng of P_{2x} cRNA and kept at 18° in physiological saline solution containing penicillin/streptomycin (11); recordings were made 2–8 days after injection. No currents in response to ATP or other agonists used in this study (0.3 or 1 mM) were recorded in uninjected oocytes.

Electrophysiological recordings. Two-electrode voltage-clamp recordings were made from oocytes using a GigaClamp amplifier (Axon Instruments); microelectrodes were filled with 3 M KCl (0.5–2 M). External solution contained 96 mM NaCl, 2 mM BaCl₂, 1 mM MgCl₂, and 0.1 mM BaCl₂; barium replacement of external calcium chloride currents (11). Conventional whole-cell recordings were made from HEK 293 cells using an Axopatch 200 patch-clamp amplifier (Axon Instruments). Patch pipettes (4–7 M Ω) contained 145 mM potassium aspartate, 11 mM EGTA, 5 mM HEPES, and 5 mM MgCl₂; external solution contained 145 mM NaCl, 2 mM KCl, 1 mM MgCl₂, 2.5 mM CaCl₂, 10 mM HEPES, and 10 mM glucose. Agonists were applied using a fast-flow U-tube delivery system (14). Native and cloned smooth muscle P_{2x} receptors from vas deferens smooth muscle express the PC-12 form of the P_{2x} receptor but were very effective agonists in oocytes and HEK cells expressing the human bladder form of the receptor (Fig. 2). For either form of the receptor, adenosine, AMP, and UTP (100 μ M) evoked currents that were 0–6% of the maximal ATP current ($n = 3$ –5 for each agonist).

Antagonists. Low concentrations of suramin (1 or 3 μ M) produced approximately parallel, rightward shifts in the ATP concentration-response curve, but the shifts in the presence of higher concentrations were no longer parallel (Fig. 3). The antagonism by even high concentrations of suramin (30 or 100 μ M) was readily reversed within 1 min of washout for HEK cells expressing the PC-12 form of the P_{2x} receptor; in the case of the bladder smooth muscle receptor the effect had been washed out by the time the agonist could be reapplied (i.e., 10 min).

The P_{2x} purinoceptor antagonist PPADS, pyridoxal-5-phosphate, oxidized ATP, and DIDS (18–22) also inhibited ATP-evoked currents in oocytes or HEK cells expressing ei-

shapes of the concentration-response curves and the calculated EC_{50} values showed no significant differences between oocyte and HEK cell expression systems. Nevertheless, because of these resolution problems in the oocyte expression system, usually only two or three agonist concentrations were applied to any one oocyte, whereas complete curves (five or six concentrations) were obtained for each agonist in individual HEK 293 cells, for both forms of the receptor. This allowed us to obtain mean EC_{50} values and to estimate 50% confidence intervals for agonists in HEK cells (see Tables 1, 2); EC_{50} values for agonists in oocytes were obtained from the pooled data, as shown in Fig. 2a. Agonists were applied in both the superfusate and the U-tube solution that contained the agonist; antagonists were superfused for 5–10 min before agonist application. Concentration-response curves for agonists and antagonists were fit by hyperbolic functions using GraphPad software (GraphPad, San Diego, CA). All data are means \pm standard errors.

Drugs. Adenosine, AMP (sodium salt), ADP (sodium salt), ATP (magnesium salt), ATP-S (tridithiol salt), UTP (sodium salt), L- β -MeATP (trithiol salt), D- β - γ -MeATP (tridithiol salt), Ba-ATP (tetraethylammonium salt), and DIDS (disodium salt) were obtained from Sigma Chemical Co. 2MeSATP (tetrasodium salt), 2-chloro-ATP (tetrasodium salt), and L- β - γ -MeATP were from Research Biochemicals International. Pyridoxal-5-phosphate monohydrate was obtained from Aldrich; APSA (tridithiol salt) was from Boehringer Mannheim, and PPADS and suramin were obtained from Bayer.

Results

Agonists. Currents evoked in responses to ATP and other purinoceptor agonists in oocytes or HEK 293 cells expressing the P_{2x} receptor desensitized in the smooth muscle form of the P_{2x} receptor, whereas the PC-12 form did not (Fig. 1; see also Refs. 11 and 12). The two expression systems yielded similar agonist concentrations, response curves and EC_{50} values (Fig. 2, Table 1). For both forms of the receptor, 2MeSATP, 2-chloro-ATP, and ADP were full agonists, whereas BaATP, AP5A, and ATP-S produced maximal responses that were about 85% of the maximal response to ATP (Fig. 2). Half-maximal concentrations (EC_{50}) values for each of these agonists to activate the PC-12 form of the P_{2x} receptor were approximately 10-fold greater than those for the human bladder form of the receptor (Table 1). The methylxylene-substituted ATP analogues α - β -MeATP, D- β - γ -MeATP, and L- β - γ -MeATP evoked little (<10%) maximal ATP current or no current in oocytes or HEK 293 cells expressing the PC-12 form of the P_{2x} receptor but were very effective agonists in oocytes and HEK cells expressing the human bladder form of the receptor (Fig. 2). For either form of the receptor, because rundown of the response is much less marked (Ref. 16 and this study), reproducible responses were obtained by applying agonist at intervals of 10 min during oocyte recordings or for 2 sec every 4 min during HEK cell recording. Little or no desensitization of the PC-12 form of the P_{2x} receptor occurs (2, 12, 17) (see Results); in these experiments, agonists were applied for similar durations but at intervals of 30–60 sec.

Agonist concentration-response curves were constructed by expressing currents as percentages of the maximal current evoked by its analogues (typically 30 or 100 μ M) in the same oocyte or HEK cell; 100 μ M was readily reversed within 1 min of washout for HEK cells expressing the PC-12 form of the P_{2x} receptor; in the case of the bladder smooth muscle receptor the effect had been washed out by the time the agonist could be reapplied (i.e., 10 min).

The P_{2x} purinoceptor antagonist PPADS, pyridoxal-5-phosphate, oxidized ATP, and DIDS (18–22) also inhibited ATP-evoked currents in oocytes or HEK cells expressing ei-

Abbreviations: α - β -MeATP, ATP- β -methylbenzoate; ATP- γ -MeATP, 3',5'-triphosphate-ATP; DIDS, 4,4'-disulfonylcyanatoaniline-2',2'-disulfone; DiS, 2-meSATP, 2-methylthio-ATP; L- β - γ -MeATP, β -methylbenzoate-ATP; AP5A, P_i-pyridoxal-5'-disulfone-5'; PPADS, pyridoxal phosphate-6-zarophenyl-2',2'-disulfone; HEK, human embryonic kidney; HEPES, 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid; EGTA, ethylene glycol bis(ether-N,N,N',N'-tetraacetic acid).

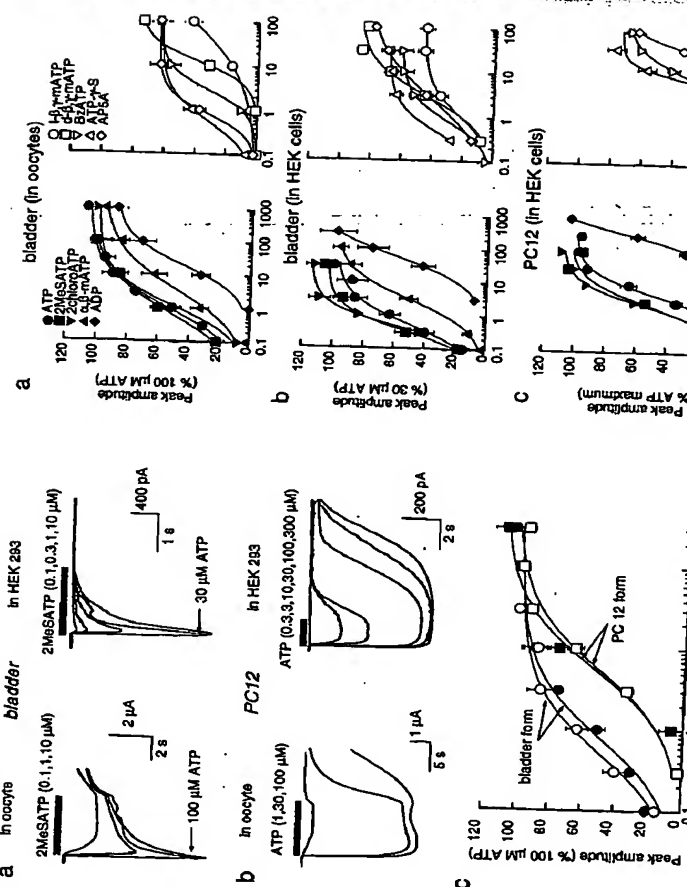


Fig. 1. Inward currents in response to activation of heterologously expressed P_{xx} purinoreceptors. **a**, Responses to activation of the P_{xx} receptor cloned from human bladder smooth muscle, expressed in HEK 293 cells (left) or HEK 293 cells (right). Superimposed current responses to increasing concentrations of ATP are shown. Note the pronounced desensitization of the smooth muscle form of the P_{xx} receptor and the absence of desensitization with the PC-12 form of the receptor. **b**, Concentration-response curves for ATP obtained with the bladder (circles) and PC-12 (squares) forms of the receptor expressed in HEK 293 cells (filled symbols) and in oocytes (open symbols). Each value is the mean \pm standard error of four to nine experiments. **c**, Concentration-response curves for ATP obtained from the rat PC-12 cell line. Superimposed currents in response to increasing concentrations of ATP are shown. Note the clear noncompetitive and required more than 15–20 min for effects to reverse (data not shown). Therefore, we measured the inhibition of the current in response to a fixed concentration of ATP (EC₅₀ concentration) with increasing concentrations of antagonist to obtain antagonist IC₅₀ values (Fig. 4). There were no clear differences in the actions of uratam, PPADS, pyridoxal-5-phosphate, or oxidized ATP to inhibit currents evoked by activation of either of the P_{xx} receptor forms; IC₅₀ values for uratam and PPADS were approximately 1–5 μ M for both receptor types and 10–20 μ M for pyridoxal-5-phosphate (Fig. 4). Oxidized ATP produced only partial inhibition of P_{xx}-mediated currents (60% inhibition at the highest concentration examined) (Fig. 4). The inhibition by oxidized ATP was reversible within 15–20 min after washout, which is in contrast to its reported action as an irreversible receptor antagonist at the "non-forming" P_{xx} purinoreceptor (22). However, this distinction may simply reflect the different antagonist incubation procedures, in that Wiley et al. (22) observed covalent linking of oxidized ATP to P_{xx} purinoreceptors in lymphocytes after 24 h of antagonist application.

Curare has been reported to be somewhat effective in blocking the P_{xx} current both in native PC-12 cells and oocytes expressing the PC-12 form of the receptor (12, 23). However, we found that only very high concentrations of curare (1 mM) produced significant inhibition of the ATP-evoked current (Fig. 4); therefore, it is unlikely that curare produced any significant inhibition of the ATP-evoked current in HEK 293 cells, with an IC₅₀ value of about 3 μ M. However, maximal concentrations of DIDS (100 and 300 μ M) produced

TABLE 1
Agonist EC₅₀ Values

Half-maximal concentrations (EC₅₀) values were calculated as the concentration giving 50% of the maximal response for each concentration-response curve generated in individual HEK 293 cells. Values are means \pm standard errors for the numbers of individual experiments shown in parentheses.

	Human urinary bladder	Rat PC-12
ATP	0.9 \pm 0.2 (6)	7.7 \pm 1 (9)
2MeSATP	0.5 \pm 0.2 (5)	2.5 \pm 1.1 (5)
2-Chloro-ATP	7.3 \pm 1.6 (5)	22.0 \pm 4.1 (5)
ADP	2.5 \pm 1.2 (5)	8.9 \pm 7 (5)
AP5A	0.7 \pm 0.2 (5)	2.3 \pm 3 (6)
BATP	3.1 \pm 0.3 (4)	2.1 \pm 3 (6)
α , β -MeATP	2.2 \pm 0.3 (5)	1.0 \pm 0.9 (7)
D ₅ , γ -MeATP	2.8 \pm 0.8 (6)	1.0 \pm 0.7 (6)
L, β , γ -MeATP	1.9 \pm 0.8 (5)	1.0 \pm 0.6 (6)

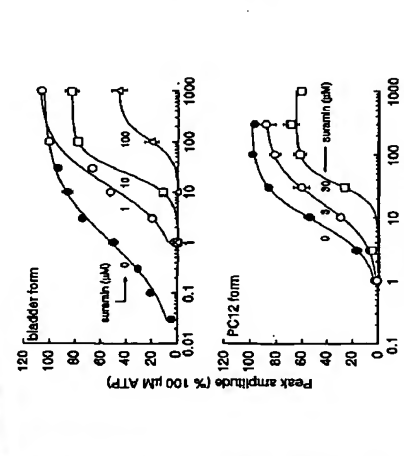


Fig. 2. Concentration-response curves for purinoreceptor agonists in the bladder smooth muscle (a and b) and PC-12 (c) forms of the P_{xx} purinoreceptor. Each value is the mean \pm standard error of four to nine separate experiments of the type illustrated in Fig. 1. **a**, Responses to activation of P_{xx} purinoreceptors in the bladder smooth muscle (filled symbols) and PC-12 form (open symbols) [ATP, 2MeSATP, 2-Chloro-ATP, ADP, AP5A for all, and α , β -MeATP (α , β -MeATP) for smooth muscle] are best fits to hyperbolic functions. Hill slopes for these lines ranged from 0.91 to 1.3. Right-hand curves for ineffective agonists are also best fits to hyperbolic functions. Note the ineffectiveness of methylxanthine analogues on the PC-12 form of the receptor. EC₅₀ values for the bladder form expressed in oocytes (a) obtained directly from these curves, are as follows: ATP, 0.8 μ M; 2MeSATP, 0.6 μ M; 2-Chloro-ATP, 0.8 μ M; ADP, 2 μ M; α , β -MeATP, 1 μ M; L, β , γ -MeATP, 2 μ M. See Table 1 for additional details of values obtained in individual cells using the HEK expression system.

TABLE 1
Agonist EC₅₀ Values

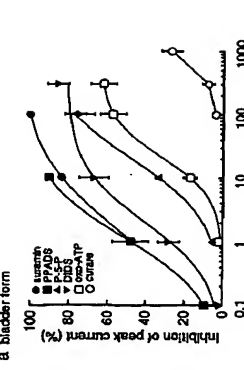


Fig. 3. Actions of suramin at the smooth muscle and PC-12 forms of the P_{xx} purinoreceptor. Each panel shows concentration-response curves for ATP in the presence of suramin. **a**, Data for the bladder form of the P_{xx} receptor, obtained in the oocyte expression system (four determinations for each value), lower, data for the PC-12 form of the receptor, obtained in the HEK cell expression system (eight determinations for values obtained in the absence of suramin and four determinations for values obtained in the presence of suramin).

<40% inhibition of the current evoked by ATP in cells expressing the PC-12 form of the receptor (Fig. 4). The original classification of the P_{xx} purinoreceptor subtype was based on the relative potency of ATP, by P_{xx} purinoreceptor antagonists produced by a fixed concentration of ATP, by P_{xx} purinoreceptor antagonists at cloned P_{xx} purinoreceptors. **b**, Inhibition of current induced by ATP at the bladder smooth muscle form of the P_{xx} receptor. Data for suramin, PPADS, and pyridoxal-5-phosphate were obtained in the oocyte expression system, whereas data for DIDS, oxo-ATP, and curare were obtained in HEK 293 cells four to six determinations for all values. **c**, Antagonism by the same antagonists of current induced by 30 μ M ATP with the PC-12 form. All data were obtained from HEK cells transiently transfected with the PC-12 P_{xx} receptor (four to eight terminations for all values).

The original classification of the P_{xx} purinoreceptor subtype was based on the relative potency of ATP and several structural analogues in assays of contractile or depolarizing actions on isolated whole tissues. This led to an agonist potency order of α , β -MeATP \gg 2MeATP \geq ATP becoming accepted as the general pharmacological definition of this receptor type (24). Although numerous subsequent studies on intact multicellular tissues yielded similar results, it has now become clear that such results were predominantly due to breakdown of ATP, 2MeATP, and other hydrolyzable analogues by ectonucleotidases. When this activity is prevented the actions of α , β -MeATP remain unaltered, whereas ATP and 2MeATP become effective at 10–100-fold lower concentrations, thus changing agonist potency to 2MeATP \geq ATP \geq α , β -MeATP. The EC₅₀ values for these agonists are all in the low micromolar (1–10 μ M) range (26) (also see Ref. 4). Whole-cell recordings from dissociated smooth muscle cells, nodose neurons, and colic neurons give similar findings (5, 6, 10), supporting the conclusion that these agonist affinities and the rank order of potencies are characteristic of endogenous smooth muscle P_{xx} purinoreceptors as well as some neuronal P_{xx} receptors. The present results on the

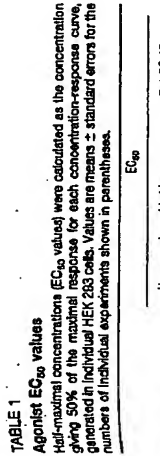


Fig. 4. Concentration-response curves for inhibition of current produced by a fixed concentration of ATP, by P_{xx} purinoreceptor antagonists at cloned P_{xx} purinoreceptors. **a**, Inhibition of current induced by 10 μ M ATP at the bladder smooth muscle form of the P_{xx} receptor. Data for suramin, PPADS, and pyridoxal-5-phosphate were obtained in the oocyte expression system, whereas data for DIDS, oxo-ATP, and curare were obtained in HEK 293 cells four to six determinations for all values. **b**, Antagonism by the same antagonists of current induced by 30 μ M ATP with the PC-12 form. All data were obtained from HEK cells transiently transfected with the PC-12 P_{xx} receptor (four to eight terminations for all values).

Discussion

The original classification of the P_{xx} purinoreceptor subtype was based on the relative potency of ATP and several structural analogues in assays of contractile or depolarizing actions on isolated whole tissues. This led to an agonist potency order of α , β -MeATP \gg 2MeATP \geq ATP becoming accepted as the general pharmacological definition of this receptor type (24). Although numerous subsequent studies on intact multicellular tissues yielded similar results, it has now become clear that such results were predominantly due to breakdown of ATP, 2MeATP, and other hydrolyzable analogues by ectonucleotidases. When this activity is prevented the actions of α , β -MeATP remain unaltered, whereas ATP and 2MeATP become effective at 10–100-fold lower concentrations, thus changing agonist potency to 2MeATP \geq ATP \geq α , β -MeATP. The EC₅₀ values for these agonists are all in the low micromolar (1–10 μ M) range (26) (also see Ref. 4). Whole-cell recordings from dissociated smooth muscle cells, nodose neurons, and colic neurons give similar findings (5, 6, 10), supporting the conclusion that these agonist affinities and the rank order of potencies are characteristic of endogenous smooth muscle P_{xx} purinoreceptors as well as some neuronal P_{xx} receptors. The present results on the

heterologously expressed P_{2X} purinoreceptor from human bladder smooth muscle directly confirm this pharmacological profile for smooth muscle P_{2X} purinoreceptors.

The present study allows direct comparisons between homodimeric forms of the smooth muscle P_{2X} receptor and the PC-12 form of this receptor, as well as between the *Xenopus* oocyte and mammalian HEK 293 cell expression systems, which are currently the two most commonly used expression systems for studying ligand-gated ion channels. We have found no obvious differences in responses of P_{2X} purinoreceptors expressed in either oocytes or HEK 293 cells (e.g., Figs. 1 and 2 and Table 1); such similarities suggest that additional proteins contributed by one or the other cell type used may not be critical for agonist/antagonist recognition. We might, therefore ask the following: what are the distinguishing features of the two receptor types with respect to agonist and antagonist binding, and how do these properties in heterologous cells compare with their properties in native cells?

We next ask how closely the properties of the expressed receptors resemble those found in the tissues from which the receptors were cloned, because discrepancies might indicate the presence of ancillary proteins that contribute to the native receptor properties. Purinoreceptor responses in rat, guinea pig, and human bladder smooth muscle have been characterized (16, 27–29); PC-12 cells have been studied by Nakazawa et al. (17, 23, 30). The properties of the human bladder form of the receptor expressed in oocytes or HEK 293 cells very closely resemble those of the native receptors; this is true with respect to the absolute concentrations of agonists that are differential sensitivity to α,β-MeATP of native smooth muscle and PC-12 P_{2X} purinoreceptors is well documented (see the introduction) and was noted in the original reports on the cloning and expression of the rat vas deferens form and the PC-12 form of these purinoreceptors (11, 12). This difference allows distinction between the smooth muscle form of the P_{2X} receptor and one kind of neuronal type of this purinoreceptor but does not allow distinction between the smooth muscle receptor and the α,β-MeATP-sensitive neuronal P_{2X} receptor (2, 4, 10). L-β,γ-MeATP may prove to be more useful for differentiation of smooth muscle P_{2X} receptors from all forms of neuronal P_{2X} receptors, because this agonist has been found to be ineffective in those neurons that do respond to α,β-MeATP (e.g., celiac and nodose ganglia), as well as those that do not (e.g., PC-12 cells and rat superior cervical ganglion neurons) (26). All other agonists examined in the present study showed approximately similar rank orders of potency for the P_{2X} receptor subtypes, although the EC₅₀ value for any one agonist to activate the PC-12 form of the receptor was approximately 10-fold greater than that for the bladder smooth muscle form (Table 1).

Of the P₂ purinoreceptor antagonists used in this study, only DIDS was able to differentiate between the smooth muscle and PC-12 forms of the P_{2X} receptor (Fig. 4). This finding may be of practical use for further delineation of P_{2X} receptor subtypes in single isolated cells, and it will be interesting to determine whether the differential sensitivity observed in the present study will be maintained at native receptors in dissociated smooth muscle and autonomic and central neurons. However, the noncompetitive and virtually irreversible nature of this inhibition at the smooth muscle P_{2X} receptor, the documented inhibition by DIDS of both P₂-like and P_{2X}-like purinoreceptors (21), and the more commonly known role of DIDS as an ion transport inhibitor are likely to limit the usefulness of DIDS as a more general P_{2X} receptor antagonist. The IC₅₀ values for suramin and PPADS to inhibit currents in response to activation of either P_{2X} receptor ranged from 1

Acknowledgments

We thank A. Brake and D. Julius for kindly providing cDNA clones for cell cultures.

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